TABLE 3—MEDICAL DEVICES FOR POSSIBLE INCLUSION IN SCOPE OF PRODUCT COVERAGE DURING OPERATIONAL PERIOD 1—Continued

Product Family	21 CFR Section No	Device Name	Tier
	892.5300	Medical neutron radiation therapy system	2
	892.5700	Remote controlled radionuclide applicator system	2
	892.5710	Radiation therapy beam-shaping block	2
	892.5730	Radionuclide brachytherapy source	2
	892.5750	Radionuclide radiation therapy system	2
	892.5770	Powered radiation therapy patient support assembly	2
	892.5840	Radiation therapy simulation system	2
	892.5930	Therapeutic x-ray tube housing assembly	1
Nuclear Medicine	892.1170	Bone densitometer	2
	892.1200	Emission computed tomography system	2
	892.1310	Nuclear tomography system	1
	892.1390	Radionuclide rebreathing system	2
General/Plastic Surgery Panel	002.1000	Thad on do not read may by other minimum.	_
Surgical Lamps	878.4630	Ultraviolet lamp for dermatologic disorders	2
Cargical Zampo	890.5500	Infrared lamp	2
	878.4580	Surgical lamp	2
Electrosurgical Cutting Equip-	878.4810	Laser surgical instrument for use in general and plastic sur-	2
ment.	070.4010	gery and in dermatology.	_
	878.4400	Electrosurgical cutting and coagulation device and accessories.	2
Miscellaneous	878.4780	Powered suction pump	2

¹Descriptive information on product codes, panel codes, and other medical device identifiers may be viewed on FDA's Internet Web Site at http://www.fda.gov/cdrh/prodcode.html.

[63 FR 60141, Nov. 6, 1998; 64 FR 16348, Apr. 5, 1999]

APPENDICES C-F TO SUBPART B OF PART 26 [RESERVED]

Subpart C—"Framework" Provisions

§ 26.60 Definitions.

- (a) The following terms and definitions shall apply to this subpart only:
- (1) Designating Authority means a body with power to designate, monitor, suspend, remove suspension of, or withdraw conformity assessment bodies as specified under this part.
- (2) Designation means the identification by a designating authority of a conformity assessment body to perform conformity assessment procedures under this part.
- (3) Regulatory Authority means a government agency or entity that exercises a legal right to control the use or sale of products within a party's jurisdiction and may take enforcement action to ensure that products marketed within its jurisdiction comply with legal requirements.
- (b) Other terms concerning conformity assessment used in this part shall have the meaning given elsewhere in this part or in the definitions contained in "Guide 2: Standardization

and Related Activities-General Vocabulary of the International Organization for Standardization (ISO) and the International Electrotechnical Commission (IEC)" (ISO/IEC Guide 2) (1996 edition), which is incorporated by reference in accordance with 5 U.S.C. 552(a) and 1 CFR part 51. Copies are available from the International Organization for Standardization, 1, rue de Varembé, Case postale 56, CH-1211 Genève 20, Switzerland, or on the Internet at http://www.iso.ch or may be examined at the Food and Drug Administration's Medical Library, 5600 Fishers Lane, rm. 11B-40, Rockville, MD 20857, or at the National Archives and Records Administration (NARA). For information on the availability of this material at NARA, call 202-741-6030, or to: http://www.archives.gov/ federal register/

code_of_federal_regulations/
ibr_locations.html. In the event of an
inconsistency between the ISO/IEC
Guide 2 and definitions in this part, the
definitions in this part shall prevail.

§26.61 Purpose of this part.

This part specifies the conditions by which each party will accept or recognize results of conformity assessment